INTRODUCTION

Intercostal schwannomas are uncommon, encapsulated neoplasms that originate in nerve sheaths of intercostal nerves. They account for less than 10% of primary neural tumors of the chest wall. Herein, we report a pathologically confirmed case of intercostal schwannoma with typical magnetic resonance imaging findings.

CASE REPORT

A 61-year-old female, having a nonspecific past history, presented to pulmonology department of our hospital with cough and sputum for about 10 days. Physical examination by a pulmonologist showed clear breathing sound. Results of her laboratory evaluation and pulmonary function test were unremarkable. However, on chest radiograph, there was nodular lesion abutting on right upper chest wall (Fig. 1A). A contrast-enhanced computed tomography (CT) scan was performed for further evaluation of this lesion. On pre-enhanced CT, about 1.5 cm sized, well-circumscribed homogenous mass was noted in right chest wall, just inferior to the 5th rib, and the mean attenuation of the lesion was 3 Hounsfield unit (HU). On CT scans acquired after contrast material administration, the mean attenuation of mass was increased to 28.7 HU and it showed peripheral enhancement (Fig. 1B). The lesion-lung interface was smooth, but it was hard to distinguish whether the mass was lo-
cated at extrapleural space or not. These CT findings of the lesion suggested a benign soft tissue tumor but could not specify the diagnosis, so we recommended MRI scan. On MRI, the mass was clearly on the lateral side of extrapleural fat which showed as thin high signal intensity lining on T1- and T2-weighted images, so it proved to be an extrapleural tumor that originated from intercostal soft tissue along the course of 5th intercostal nerve in the right side of chest wall. The mass showed low signal intensity (similar to that of adjacent muscles) on T1-weighted MR images, and intermediate to high signal intensity with in-

Fig. 1. A 61-year-old woman with intercostal schwannoma involving the right 5th intercostal nerve.
A. Chest radiography shows a nodular lesion abutting on right upper chest wall, just inferior to 5th rib (arrows).
B. Pre-enhanced CT demonstrates a 1.5 cm nodule showing heterogeneous attenuation in the chest wall, just inferior to the 5th rib. After CE, the mass shows thin peripheral enhancement.
C. The mass reveals low signal intensity on T1WI, intermediate signal intensity with internal high signal intensity focus (arrowhead) on T2WI. On gadolinium-enhanced fat-suppressed T1WI, it shows bright enhancement, with tiny non-enhancing area (arrowhead) corresponding to the high signal intensity focus on T2WI.
D. Intraoperative thoracoscopic image demonstrates a well circumscribed mass abutting on inferior margin of the right 5th rib showing mass effect on the chest wall.
CE = contrast enhancement, CT = computed tomography, T1WI = T1-weighted image, T2WI = T2-weighted image
ternal high signal intensity focus on T2-weighted images. On gadolinium-enhanced fat-suppressed T1-weighted images, it showed bright enhancement with small non-enhancing area that was consistent with high signal intensity focus on T2-weighted image, suggesting central cystic degeneration (Fig. 1C). Based on these MRI findings, the radiologic differential diagnosis was assumed as neurogenic tumor that originated in right 5th intercostal nerve. The patient underwent mass excision via video-assisted thoracoscopic surgery. The mass was 1.5 cm sized and located at inferior aspect of right 5th rib, abutting superficial surface of parietal pleura (Fig. 1D). There was no remarkable bony erosion on right 5th rib. The final pathologic diagnosis was intercostal schwannoma (Fig. 1E, F).

**DISCUSSION**

Schwannomas are type of benign peripheral nerve sheath neoplasm. The tumor has a propensity to involve head, neck, flexor surfaces of extremities, posterior mediastinum, and retroperitoneum (6). Chest wall is an uncommon location for schwannomas and they arise from spinal nerve roots or intercostal nerves (7), but they may involve any thoracic nerves (including the phrenic or vagus nerve) (5). The incidence of intercostal schwannomas is below 10% of all primary neurogenic thoracic tumors (1). Schwannomas typically occur in patients between 20 years and 50 years of age, with no sex predilection (2). They are almost slow-growing nonaggressive neoplasms and usually manifest clinically as a painless mass without neurologic symptoms, unless the mass has become large enough to compress the adjacent nerve (4, 8).

Grossly, small tumors tend to be spheroid, firm, and well circumscribed, whereas larger tumors are ovoid or irregularly lobulated (2). Chest radiographs do not usually depict small schwannomas, but when the lesion is delineated, it shows a smoothly margined, oval mass (2, 5). Unenhanced CT scans of schwannoma show a well-circumscribed homogeneous mass with attenuation slightly less than or equal to that of muscle (2). Schwannomas may contain areas of low attenuation corresponding to fat or cystic degeneration, and calcifications may be primarily in a peripheral pattern, particularly in long-standing lesions with advanced degeneration (8). Adjacent osseous pressure erosion may also be present, but it is relatively rare, and presence of bone erosion without destruction indicates the benign nature and slow growth rate of the lesion (2, 5). On contrast-enhanced CT, the attenuation of the mass is equal to or slightly greater than that of muscle (2). When the mass is small, the mass may show homogeneous enhancement, while larger schwannomas can contain nonenhancing area and show more heterogeneous enhancement due to cystic and hemorrhagic changes (2, 5).

MRI shows low to intermediate signal intensity (equal to or slightly greater than that of muscle) on T1-weighted images, and shows relatively high signal intensity on T2-weighted images. On gadolinium-enhanced imaging, schwannomas show intense enhancement of solid components (2, 5). The larger the schwannoma...
MRI Finding of Intercostal Schwannoma

noma, the more heterogeneous it will appear on all sequences due to cystic degeneration, hemorrhage, or both. Regions of very high signal intensity on T2-weighted imaging correspond to cystic degeneration and may only show peripheral zone enhancement or no enhancement associated with the very high signal intensity, and those findings favor schwannoma (5). Several foci of marked hyperintensity on T2-weighted image due to multifocal cystic degeneration, which is so-called bead-like appearance, are one of the characteristic MRI finding of schwannoma (9). Also, as in our case, MRI can clearly demonstrate extrapleural fat which shows high signal intensity on T1- and T2-weighted images, so it can help distinguish extrapleural tumors from pleural tumors.

These imaging findings of schwannomas are similar to those seen with neurofibromas and, in many cases, cannot be distinguished (8), so fine-needle cytology or surgical biopsy is often needed to establish the diagnosis (10). However, some features can help differentiate these two lesions. Pathologically, schwannomas are fusiform masses that are eccentrically located in relation to the involved nerve, so in imaging studies, schwannomas tend to be eccentrically positioned and the parent nerve can be identified, whereas centrally located masses suggest neurofibromas. In addition, heterogeneous appearance with degeneration and cystic cavitation are much more common in schwannomas than in neurofibromas (8). Areas designated as Antoni A are more organized and are composed of cellular spindle cells arranged in short bundles or interlacing fascicles. Antoni B areas are hypocellular and contain more myxomatous loosely arranged tissue (7). In our case, upper margin of the mass was just abutting on the lower margin of the rib where the intercostal nerve located, and extended mainly to the inferior direction, resulting eccentric position related to intercostal nerve. And cystic degeneration was also noted within the mass. From these imaging features, the diagnosis of our case could be intercostal schwannoma more likely, rather than neurofibroma.

Besides neurofibroma, there are other thoracic neurogenic tumors that should be considered as differential diagnosis, including neuroblastoma, paraganglioma, ganglioneuroma, and malignant peripheral nerve sheath tumor. There are some imaging features that can help differentiate these tumors from schwannoma. Neuroblastomas can show coarse or curvilinear calcifications on CT. Paragangliomas show marked enhancement due to increased vascularity and may show salt-and-pepper appearance from vessel signal flow voids within the mass on T2-weighted MR images. Ganglioneuromas are usually elongated and oriented on a vertical axis following the direction of the sympathetic chain, and may show whorled appearance of low signal intensity bands on T1- and T2-weighted MR images. And, malignant peripheral nerve sheath tumors (MPNSTs) usually show compression or destruction of adjacent structures and pleural abnormalities indicating malignancy. Some MRI features are useful in diagnosis of MPNSTs, and if a tumor has two or more features, it should be considered highly suspicious of malignancy. Those features are increased largest dimension of the mass (> 5 cm), presence of peripheral enhanced pattern, presence of perilesional edema, and presence of intratumoral cystic change. In our case, though the mass showed tiny focus of cystic change, there was no other features that suggesting malignancy, so it is more likely to be benign (5).

Local resection of the mass is sufficient treatment for most of the smaller schwannomas of intercostal nerve, and thoracoscopy is usually the preferred surgical approach. Larger or more aggressive tumors require resection of the chest wall (3). The possibility of recurrence or malignant transformation is considered extremely low (10).

In summary, we present a case of schwannoma of intercostal nerve that shows obvious and typical imaging features, especially on MRI. Although pathologic confirm is recommended for diagnostic confirmation of this entity, characteristic MRI features may provide a clue to the diagnosis and play an important role in the diagnosis of schwannoma.

REFERENCES

늑간 신경초종의 MRI 소견: 증례 보고

심하얀1 · 양 익1 · 홍혜숙1 · 우지영1 · 황지영1 · 문진희1 · 김한면1 · 김혜정2 · 황숙민1 · 신미경2 · 김희영1*

늑간 신경초종은 늑간 신경에서 기원하는 피막으로 둘러싸인 드문 종양이며, 흉곽에서 발생하는 신경성 종양의 10% 미만을 차지한다. 우리는 특징적인 자기공명영상 소견을 보이는 병리학적으로 확진된 늑간 신경초종에 대한 증례를 보고한다.

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